WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁵ :		11) International Publication Number: WO 92/2228
A61K 9/72, C09K 3/30	A1	43) International Publication Date: 23 December 1992 (23.12.92
(21) International Application Number: PCT/US (22) International Filing Date: 8 June 1992		poration, One Giralda Farms, Madison, NJ 07940-100
(30) Priority data: 712,791 10 June 1991 (10.06.91) (60) Parent Application or Grant (63) Related by Continuation US Filed on 10 June 1991 (71) Applicant (for all designated States except US): SC CORPORATION [US/US]; 2000 Galloping I Kenilworth, NJ 07033 (US). (72) Inventors; and (75) Inventors/Applicants (for US only): FASSBERG [US/US]; 175 W. 12th Street, New York, NY 10 SEQUEIRA, Joel, A. [US/US]; 18 Stuyvesant York, NY 1009 (US). CHAUDRY, Imitaz, A. 18 Rose Avenue, North Caldwell, NJ 07 KOPCHA, Michael [US/US]: 141 Wycoff V. East Brunswick, NJ 08816 (US).	,791 (C. (10.06.9) (HERIN Hill Rose (), Julian (0011 (U Oval, N (US/U (006 (U	patent), GR (European patent), HU, IT (European patent), HV, RF, KR, LK, LU (European patent), MC (European patent), MC, GR, CE, CE, CE, CE, CE, CE, CE, CE, CE, CE

(57) Abstract

Aerosol formulations substantially free of chlorofluorocarbons for oral and/or nasal administration are described. The formulations comprise 1,1,1,2,3,3,3 heptafluoropropane, a medicament, optionally an excipient and optionally a surfactant. Methods of treatment utilizing the formulations are also described.

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NON-CHLOROFLUOROCARBON AEROSOL FORMULATIONS

INTRODUCTION TO THE INVENTION

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The present invention is directed at aerosol formulations which are substantially free of chlorofluorocarbons (CFC's). More specifically, the present invention is directed at formulations substantially free of CFC's and having particular utility in medicinal applications, especially in metered dose pressurized inhalators (MDI's).

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Metered dose inhalators have proven to be an effective method for delivering medicaments orally and nasally. They have been used extensively for delivering bronchodilating and steroidal compounds to asthmatics and may also be useful for delivering other compounds such as pentamidine and non-bronchodilator anti-inflammatory drugs. The rapid onset of activity of compounds administered in this manner and the absence of any significant side effects have resulted in a large number of compounds being formulated for administration via this route. Typically, the drug is delivered to the patient by a propellant system generally comprising one or more propellants which have the appropriate vapor pressure and which are suitable for oral or nasal administration. The more preferred propellant systems typically comprise propellant 11, propellant 12, propellant 114 or mixtures thereof. Often the vapor pressure of the propellant systems is adjusted by admixing a liquid excipient with the propellant.

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However, propellants 11, 12 and 114 belong to a class of compounds known as chlorofluorocarbons, which have been linked to the depletion of ozone in the atmosphere. It has been postulated that

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ozone blocks certain harmful UV rays and that a decrease in the atmospheric ozone content will result in an increase in the incidence of skin cancer. In the 1970's certain steps were taken to reduce the CFC emissions from aerosols. Other propellants, such as hydrocarbons, were used, or the product was delivered in a different manner. Because CFC usage in medicinal applications is relatively low i.e. less than 1% of total CFC emissions, and because of the health benefits associated with metered dose inhalators, steps were not taken at that time to restrict the use of CFC propellants in metered dose inhalators.

However, continuing and more sophisticated ozone measurements have indicated that the earlier restrictions in CFC usage were insufficient and that additional, significant steps should be taken to drastically reduce CFC emissions. Recently, recommendations have been made that CFC production be virtually discontinued by the end of this century. As a result, it may not be possible to continue to use CFC propellants in the intermediate and long term. While some efforts have been made to use non-pressurized metered dose inhalators, many of these devices have not been completely successful. Many do not deliver uniform doses, are mechanically complex, do not provide the 100-200 doses per unit of current aerosol containers, are difficult for individuals to utilize, and are bulky and/or cumbersome for the patients to use, particularly when they have an acute need for the medication.

As a result, there is a need for aerosol formulations which are substantially free of CFC's. Non-CFC propellants systems must meet several criteria for pressurized metered dose inhalators. They must be non-toxic, stable and non-reactive with the medicament and the other major components in the valve/actuator. One propellant which has been found to be suitable is CF₃-CH₂F-CF₃, also known as Freon 227, HFA 227, HFC 227 or 1,1,1,2,3,3,3 heptafluoropropane. However, certain physical properties, i.e., polarity and solubility, of HFC 227 differ from those of commonly used CFC propellants. Commonly used surfactants may be insoluble in HFA 227. Moreover, where the medicament is to be delivered as a solution, the medicament may not be readily soluble in this propellant. The polarity difference between HFC 227 and the previously used CFC propellants may result in a different delivery of the medicament when HFC 227 replaces a CFC propellant.

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The medicament may cream, settle or agglomerate in the non-CFC propellant even though this did not occur in the CFC propellant.

The use of HFA 227 previously has been disclosed for use in medicinal inhalators. European Patent Publication No. 0 384 371 is directed at the combination of propellant 227 and propane, butane, isobutane, Me₂O and/or F₂CHMe.

Research Disclosure No. 30161, May, 1989 discloses that non-CFC propellants, such as fluorohydrocarbons may be used in pressurized medicaments delivered directly to the lungs, e.g. bronchodilators.

Other publications have been directed at the use of other fluorohydrocarbons, such as HFC 134a, for aerosol propellants. European Patent Publication No. 0 372 777 is directed at medicinal aerosol formulations incorporating HFC 134a and an adjuvant having a higher polarity than the propellant. This publication lists several possible adjuvants and surfactants for use in combination with the propellant and the medicament.

International patent application No. WO 91/04011 discloses the combination of HFC 134a and a powdered medicament pre-coated with a non-perfluorinated surfactant prior to dispersing the powdered medicament in the propellant. Pages 6-7 of the publication list suitable surfactants for use with the propellant. A perfluorinated adjuvant optionally could be added. However, the pre-coating of the medicament may not be advantageous, since it adds an additional, complex step to the manufacturing process.

U.S. Patent No. 4,174,295 discloses the combination of HFC 134a with various chlorofluorocarbons and optionally a saturated hydrocarbon. U.S. Patent No. 2,885,427 discloses the use of HFC-134a as an aerosol propellant. U.S. Patent No. 3,261,748 discloses the use of HFC-134a for anesthesia. U.S. Patent Nos. 4,129,603, 4,311,863, 4,851,595 and European Publication No. 379,793 also disclose the use of HFC-134a as an aerosol propellant.

However, the specific combinations noted above may not provide the desired solubility, stability, low toxicity, exact dosage, correct particle size (if suspension) and/or compatibility with commonly used valves assemblies of metered dose inhalers.

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SUMMARY OF THE INVENTION

Accordingly, the present invention is directed at a non-toxic formulation substantially free of CFC's having improved stability and compatibility with the medicament and which is relatively easily manufactured.

The present invention also is directed at formulations which may be utilized in present aerosol filling equipment with only relatively minor modifications and without pre-coating the medicament.

One embodiment of the present invention is directed at a formulation comprising:

A. Propellant 1,1,1,2,3,3,3 heptafluoropropane;

B. optionally an excipient selected from the group consisting of alcohols, Miglyol 812, Miglyol 840, PEG-400, menthol, lauroglycol, Vertrel 245, Transcutol, Labrafac Hydro WL 1219,

perfluorocyclobutane, eucalyptus oil, short chain fatty acids, and combinations thereof;

C. a medicament; and

group consisting of oleic acid, sorbitan trioleate, cetyl pyridinium chloride, soya lecithin, Tween 20, Tween 60, Tween 80, Pluronic L-121

and Pluronic L-92, castor oil ethoxylate, pluronic F 68, Tetronic 150 R1 and combinations thereof.

Also included within the invention is an aerosol formulation 25 comprising:

A. an effective amount of medicament;

B. 1,1,1,2,3,3,3 heptafluoropropane; and

an excipient selected from the group consisting
 of: propylene glycol diesters of medium chain fatty
 acids;

triglyceride esters of medium chain fatty acids:

perfluorodimethylcyclobutane; perfluorocyclobutane;

polyethylene glycol;

menthol;

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	lauroglycol;
	diethylene glyco! monoethylether;
	polyglycolized glycerides of medium chain
	fatty acids;
5	alcohols;
	eucalyptus oil;
	short chain fatty acids;
	and combinations thereof.

The formulation optionally may further comprise a surfactant. The surfactant preferably is selected from the group consisting of:

	oleic acid;
	sorbitan trioleate;
15	cetyl pyridinium chloride;
	soya lecithin;
et.	polyoxyethylene(20) sorbitan monolaurate;
	polyoxyethylene (10) stearyl ether;
	polyoxyethylene (2) oleyl ether;
20	polyoxypropylene-polyoxyethylene-ethylene diamine block copolymers;
	polyoxyethylene(20) sorbitan monostearate;
	polyoxyethylene(20) sorbitan monooleate;
	polyoxypropylene-polyoxyethylene block
25	copolymers;
	castor oil ethoxylate; and combinations
	thereof.

The preferred liquid excipients are diethylene glycol monethyether, propyleneglycol diesters of medium chain fatty acids, perfluorodimethylcyclobutane and polyethylene glycol.

The preferred surfactants are oleic acid; sorbitan trioleate, cetylpyridinium chloride; polyoxyethylene (20) sorbitan monolaurate; polyoxypropylene-polyoxyethylene block copolymers; soya lecithin; and polyoxypropylene-polyoxyethylene-ethylenediamine block copolymers; with oleic acid being particularly preferred.

The invention is of particular utility where the medicament is albuterol, mometasone furoate or beclomethasone dipropionate, and salts and clathrates thereof.

A useful formulation range comprises:

5	Α.	1,1,1,2,3,3,3 heptafluoropropane	25 - 99.99 wt %
	В.	medicament	0.01 - 1 wt %
	C.	excipient	0 - 75 wt %
	D	surfactant	0 - 3 wt %

The present invention also is directed at a method of treating asthma in mammals comprising administering to a mammal in need of such treatment an effective amount of aerosol formulation comprising:

A. a medicament selected from the group comprising
15 albuterol, mometasone furoate, beclomethasone dipropionate, and salts
and clathrates thereof;

B. 1,1,1,2,3,3,3 heptafluoropropane; and

C. optionally an excipient selected from the group consisting of:

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propylene glycol diesters of medium chain fatty acids;
triglyceride esters of medium chain fatty acids;
perfluorodimethylcyclobutane;
perfluorocyclobutane;
polyethylene glycol;
menthol;
lauroglycol;
diethyleneglycol monoethylether;
polyethycolized glycerides of medium chain

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polyglycolized glycerides of medium chain fatty acids;

fatty acids; alcohols:

short chain fatty acids;

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eucalyptus oil; and combinations thereof.

A surfactant optionally is present. The surfactant preferably is selected from the group consisting of:

oleic acid: 5 sorbitan trioleate: cetyl pyridinium chloride; sova lecithin: polyoxyethylene (20) sorbitan monolaurate; polyoxyethylene (10) stearyl ether: 10 polyoxyethylene (2) oleyl ether: polyoxyethylene-polyoxypropylene-ethylene diamine block copolymers: polyoxyethylene (20) sorbitan monostearate; polyoxypropylene-polyoxyethylene block 15 copolymers: castor oil ethoxylate; and combinations thereof.

DETAILED DESCRIPTION OF THE INVENTION

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The formulations of the present invention all utilize propellant 227 in combination with the medicament, optionally a liquid excipient and optionally a surfactant.

The excipient facilitates the compatibility of the medicament with the propellant and also lowers the discharge pressure to an acceptable range i.e. about $2.76 - 5.52 \times 10^5$ newton/meter² absolute (40 to 80 psia), preferably $3.45 - 4.83 \times 10^5$ newton/meter² absolute (50 to 70 psia). The excipient chosen must be non-reactive with the medicament, relatively non-toxic, and should have a vapor pressure below about 3.45×10^5 newton/meter² absolute (50 psia). As used hereinafter the term "medium chain fatty acids" refers to chains of alkyl groups terminating in a -COOH group and having 6-12 carbon atoms, preferably 8-10 carbon atoms. The term "short chain fatty acids" refers to chains of alkyl groups terminating in a -COOH group and having 4-8 carbon atoms. The term "alcohol" includes C_1 - C_3 alcohols, such as methanol, ethanol and isopropanol. Among the preferred excipients are:

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propylene glycol diesters of medium chain fatty acids available under the tradename Miglyol 840 (from Hüls America, Inc. Piscataway, N.J.);

triglyceride esters of medium chain fatty acids available under the tradename Miglyol 812 (from Hüls);

perfluorodimethylcyclobutane available under the tradename Vertrel 245 (from E.I DuPont de Nemours and Co. Inc. Wilmington, Delaware);

perfluorocyclobutane available under the tradename octafluorocyclobutane (from PCR Gainsville, Florida);

polyethylene glycol available under the tradename PEG 400 (from BASF Parsippany, N.J.);

menthol (from Pluess-Stauffer International Stanford, Connecticut);

propylene glycol monolaurate available under the tradename lauroglycol (from Gattefossé Elmsford, N.Y.);

diethylene glycol monoethylether available under the tradename Transcutol (from Gattefossé);

polyglycolized glyceride of medium chain fatty acids available under the tradename Labrafac Hydro WL 1219 (from Gattefossé):

alcohols, such as ethanol, methanol and isopropanol; eucalyptus oil available (from Pluess-Stauffer International); and mixtures thereof.

A surfactant optionally may be added to lower the surface and interfacial tension between the medicament and the propellant. Where the medicament, propellant and excipient are to form a suspension, a surfactant may or may not be required. Where the medicament, propellant and excipient are to form a solution, a surfactant may or may not be necessary, depending in part, on the solubility of the particular medicament and excipient. The surfactant may be any suitable, non-toxic compound which is non-reactive with the medicament and which substantially reduces the surface tension between the medicament, the excipient and the propellant and/or acts as a valve lubricant. Among the preferred surfactants are:

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oleic acid available under the tradename oleic acid NF6321 (from Henkel Corp. Emery Group, Cincinnati, Ohio); cetylpyridinium chloride (from Arrow Chemical, Inc. Westwood, N.J.);

soya lecithin available under the tradename Epikuron 200 (from Lucas Meyer Decatur, Illinois);

polyoxyethylene(20) sorbitan monolaurate available under the tradename Tween 20 (from ICI Specialty Chemicals, Wilmington, Delaware);

polyoxyethylene(20) sorbitan monostearate available under the tradename Tween 60 (from ICI);

polyoxyethylene(20) sorbitan monooleate available under the tradename Tween 80 (from ICI);

polyoxyethylene (10) stearyl ether available under the tradename Brij 76 (from ICI);

polyoxyethylene (2) oleyl ether available under the tradename Brij 92 (from ICI);

polyoxyethylene-polyoxypropylene-ethylenediamine block copolymer available under the tradename Tetronic 150 R1 (from BASF);

polyoxypropylene-polyoxyethylene block copolymers available under the tradenames Pluronic L-92, Pluronic L-121 and Pluronic F 68 (from BASF);

castor oil ethoxylate available under the tradename Alkasurf CO-40 (from Rhone-Poulenc Mississauga Ontario,Canada); and mixtures thereof.

The medicaments of the present invention may include any pharmaceutically active compounds which are to be delivered by oral inhalation or nasally. Typical classes of compounds include bronchodilators, anti-inflammatory compounds, antihistamines, antiallergics, analgesics, antitussives, anti-anginal medications, steroids, corticosteroids, vasoconstrictors and antibiotics. Specific compounds within these classes of compounds are albuterol, mometasone furoate, beclomethasone dipropionate, isoproterenol, heparin, terbutaline, rimiterol, perbuterol, disodium cromoglycate, isoprenaline, adrenaline, pentamidine and ipratropium bromide. These compounds may be utilized either as the free base, as a salt, or as a

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clathrate, depending upon the stability and solubility of the active compound in the specific formulation. When clathrates are utilized, P-11 and hexane clathrates are particularly preferred.

Where the active compound forms a suspension, the particle size should be relatively uniform, with substantially all the particles preferably ranging between about 0.1-25 microns, preferably 0.5-10 microns, more preferably 1-5 microns. Particles larger than 25 microns may be held up in the oropharyngeal cavity, while particles smaller than about 0.5 micron preferably are not utilized, since they would be more likely to be exhaled and, therefore, not reach the lungs of the patient.

The formulations of the present invention may be filled into the aerosol containers using conventional filling equipment. Since propellant 227 may not be compatible with all elastomeric compounds currently utilized in present aerosol valve assemblies, it may be necessary to substitute other materials, such as white buna rubber, or to utilize excipients and optionally surfactants which mitigate the adverse effects of propellant 227 on the valve components.

To assure uniform dispersion of the active ingredient, the formulations typically will include the following components:

	Range (wt %)	Preferred Range (wt%)	Most Preferred Range (wt%)
Medicament	0.01 - 1	0.03 - 0.7	0.05 - 0.5
Propellant	25 - 99.99	50 - 99.97	50 - 99.95
Excipient(s)	0 - 75	0 - 50	0 - 50
Surfactant(s)	0 - 3	0 - 2	0 - 1

Depending on the particular application, the container may be charged with a predetermined quantity of formulation for single or multiple dosing. Typically, the container is sized for multiple-dosing, and, therefore, it is very important that the formulation delivered is substantially uniform for each dosing. For example, where the formulation is for bronchodilation, the container typically is charged with a sufficient quantity of the formulation for 200 charges.

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Suitable suspensions may be screened in part by observing several physical properties of the formulation, i.e. the rate of particle agglomeration, the size of the agglomerates and the rate of particulate creaming/settling and comparing these to an acceptable standard. Suitable solutions may be screened by observing the solubility of the medicament over the entire recommended storage temperature range.

Suspensions of the present invention preferably may be prepared by either the pressure filling or cold filling procedures well-known in the art.

For metered dose inhalators, suspensions may be particularly preferred for efficacy and stability considerations.

Those skilled in the art may choose to add one or more preservative, buffer, antioxidant, sweetener and/or flavors or other taste masking agents depending upon the characteristics of the formulation.

Examples I - XXXIII below further describe the present invention. For several of the examples, alternative formulations denoted as A and B are provided.

Component Wt%

EXAMPLE I

-	A	В	
Albuteroi	0.5	0.1	
Miglyol 812	10.0	1.0	
HFC-227	89.5	98.9	

EXAMPLE II

Albuterol	0.1
Transcutol	25.0
HFC-227	74.9

EXAMPLE III

		Α	В
Albuterol	(0.5	0.1
Miglyol 840	10	0.0	1.0
HFC-227	89	9.5	98.9
E	XAMPLE IV		-
Albuterol		0.1	
PEG 400		1.0	
HFC-227	98	8.9	
E	XAMPLE V		
		0.1	
Albuterol		0.5	
Menthol		98.9	
HFC 227		00.0	
E	XAMPLE VI		
		Α	В
Albuterol	1	0.1	0.1
Laurogiycol		0.1	0.5
HFC 227	9	9.8	99.4
E	XAMPLE VII		
			В
		A	0.5
Albuterol		0.1 0.0	49.6
Vertrel 245			49.9
HFC 227	8	9.9	4 ∂.∂

EXAMPLE VIII

Albuterol Labrafac Hydro WL 1219 HFC 227		0.1 0.5 99.4	
	EXAMPLE IX		
		A	В
Albuterol		0.1	0.5
Perfluorocyclobutane		10.0	49.6
HFC 227		89.9	49.9
	EXAMPLE X		
		Α	В
Oleic Acid		0.01	0.1
Albuterol		0.10	0.1
Ethanol		1.00	30.0
HFC 227		98.89	69.8
	EXAMPLE XI		
		A	В
Oleic Acid		0.01	0.1
Albuterol sulfate		0.10	0.1
Ethanol		1.00	30.0
HFC 227		98.89	69.8
	EXAMPLE XII		
		Α	В
Oleic Acid		0.01	0.1
Albuterol		0.10	0.1
Ethanol		1.00	25.0
HFC 227		98.89	74.8

HFC-227

	EXAMPLE XIII		
		Α	В
Oleic Acid		0.001	0.01
Albuterol		0.1	0.1
Miglyol 812		1.0	10.0
HFC 227		98.8	89.8
	EXAMPLE XIV		
Tetronic 150 R1		0.1	
Albuterol		0.1	
Miglyol 812		9.8	
HFC-227		90	
AFO-221		00	
	CVALIDI E MI		
	EXAMPLE XV	Δ.	В
DI 1.1404		A 	0,1
Pluronic L121		0.1	0.1
Albuterol		1.0	10.0
Miglyol 812		98.8	89.8
HFC 227		90.0	03.0
-	EXAMPLE XVI		
Tween 20		0.1	
Albuterol		0.1	
Miglyol 812		10.0	
Vertrel 245		10.0	
·			

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EXA	ME	ᄓᆮ	XX	/11

	A_	В	
Oleic Acid	0.01	0.1	
Albuterol Sulfate	0.10	0.1	
Ethanol	1.00	25.0	
HFC 227	98.89	74.8	
	EXAMPLE XVIII	_	
Olate Astal	A	<u>B</u>	
Oleic Acid	0.01		
Albuterol Sulfate	0.10		
Transcutol	1.00	25.0	
HFC 227	98.89	74.8	
EXAMPLE XIX			
	A	B	
Pluronic L 121	0.1	0.1	
Mometasone Furoate	0.1	0.1	
Miglyol 812	1.0	10.0	
HFC 227	98.8	89.8	
EXAMPLE XX			
Tetronic 150 R1	0.1		
Mometasone Furoate	0.1		
Miglyol 812	9.8		
			

EXAMPLE XXI

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Mometasone Furoate	0.1
HFC-227	99.9

HFC-227

EXAMPLE XXII

Beclomethasone Dipropionate	0.1
HFC-227	99.9

EXAMPLE XXIII

Mometasone Furoate	0.1
Tween 20	0.01
HFC-227	99.89

EXAMPLE XXIV

Beclomethasone Dipropionate	0.1
Tween 20	0.01
HFC-227	99.89

EXAMPLE XXV

Mometasone Furoate	0.1
Tween 20	0.01
Oleic Acid	0.0005
HFC-227	99.8895

EXAMPLE XXVI

Beclomethasone Dipropionate	0.1
Tween 20	0.01
Oleic Acid	0.0005
HFC-227	99.8895

EXAMPLE XXVII

Mometasone Furoate	0.1
Miglyol 812	9
Oleic Acid	0.005
Tetronic 150 R1	0.01
HFC-227	90.885

EXAMPLE XXVIII

Beclomethasone Dipropionate	0.1
Miglyol 840	9
Oleic Acid	0.005
Pluronic L121	0.01
HFC-227	90.885

EXAMPLE XXIX

	A	B	_
Oleic Acid	0.001	0.01	
Mometasone Furoate	0.1	0.1	
Miglyol 812	1.0	10.0	
HFC 227	98.8	89.8	

EXAMPLE XXX

	A	<u>-</u> В	
Pluronic L121	~ 0.1	0.1	
Beclomethasone Dipropionate	0.1	0.1	
Miglyot 812	1.0	10.0	
HFC 227	98.8	89.8	

EXAMPLE XXXI

	A	В	
Beclomethasone Dipropionate	0.1	0.1	
Miglyol 812	1.0	10.0	
HFC 227	98.9	89.9	

EXAMPLE XXXII

_	A	В В	
Beclomethasone Dipropionate	0.1	0.1	
PEG 400	1.0	10.0	
HFC 227	98.9	89.9	

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FXAMPLE XXXIII

Beclomethasone Dipropionate	0.1
Ethanol	5
HFC 227	94.9

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While the examples above have been directed at albuterol, albuterol sulfate, mometasone furoate, beclomethasone dipropionate and beclomethasone dipropionate clathrates, it is contemplated that other orally or nasally administered medicaments could be utilized. Similarly, it is contemplated that excipients and surfactants other than those exemplified may be utilized.

The descriptions of the foregoing embodiments of the invention have been presented for the purpose of illustration and description. They are not intended to be exhaustive or to limit the invention to the precise forms disclosed, and obviously many modifications and variations are possible in light of the above teaching. The embodiments were chosen and described in order to best explain the principles of the invention and its practical application to thereby enable others skilled in the art to best utilize the invention in various embodiments and with various modifications as are suited to the particular use contemplated. It is intended that the scope of the invention be defined by the claims appended hereto.

What is claimed is:

1. An aerosol formulation consisting essentially of:

5 A. an effective amount of a medicament;

B. 1,1,1,2,3,3,3 heptafluoropropane; and optionally, one or more components selected from one or more of the following:

excipients;

10 surfactants; and

additives which are:

preservatives;

buffers;

antioxidants;

15 sweetners; and

taste masking agents.

2. The formulation of claim 1 wherein the excipient is selected from the group consisting of:

20 propylene glycol diesters of medium chain fatty acids;

triglyceride esters of medium chain fatty acids:

perfluorodimethylcyclobutane;

perfluorocyclobutane;

polyethylene glycol;

25 menthol:

laurogiycol;

diethylglycol monoethylether;

polyglycolized glycerides of medium chain fatty acids;

alcohols:

30 short chain fatty acids;

eucalyptus oil; and combinations thereof.

3. The formulation of claim 1 wherein the surfactant is selected from the group consisting of:

35 oleic acid:

sorbitan trioleate;

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cetyl pyridinium chloride;
soya lecithin;
polyoxyethylene (20) sorbitan monolaurate;
polyoxyethylene(20) sorbitan monostearate;
polyoxyethylene(20) sorbitan monooleate;
polyoxyethylene (10) stearyl ether;
polyoxyethylene (2) oleyl ether;
polyoxyethylene-polyoxypropylene-ethylenediamine block
copolymers;
polyoxypropylene-polyoxyethylene block copolymers;
castor oil ethoxylate; and combinations thereof.

- The formulation of claim 1 wherein the medicament is selected from the group consisting of: albuterol, mometasone furoate,
 beclomethasone dipropionate, isoproterenol, heparin, terbutaline, rimiterol, perbuterol, disodium cromoglycate, isoprenaline, adrenaline, pentamidine, ipratropium bromide, and salts and clathrates thereof.
 - 5. The formulation of claim 4 wherein the medicament is selected from the group consisting of:

albuterol, albuterol sulfate, beclomethasone dipropionate, beclomethasone dipropionate clathrates and mometasone furoate.

6. The formulation of claim 5 which is substantially free of chlorofluorocarbons.

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7. The formulation of claim 5 containing an excipient selected from the group consisting of diethylene glycol monoethylether, propylene glycol diesters of medium chain fatty acids, perfluorodimethylcyclobutane and polyethylene glycol.

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8. The formulation of claim 7 containing a surfactant selected from the group consisting of: oleic acid, sorbitan trioleate, cetylpyridinium chloride and soya lecithin.

9. The formulation of claim 1 containing the following components in the indicated ranges:

	medicament	0.01- 1 wt %
5	1,1,1,2,3,3,3 heptafluoropropane	25 - 99.99 wt %
	excipient	0 - 75 wt %
	surfactant	0 - 3 wt %

10. The formulation of claim 9 containing the following components inthe indicated ranges:

medicament	0.03 - 0.7 wt%
1,1,1,2,3,3,3 heptafluoropropane	50 - 99.97 wt%
excipient	0 - 50 wt%
surfactant	0 - 2 wt%

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11. The formulation of claim 10 containing the following components in the indicated ranges:

	medicament	0.05 - 0.5 wt%
20	1,1,1,2,3,3,3 heptafluoropropane	50 -99.95 wt%
	excipient	0 - 50 wt%
	surfactant	0 - 1 wt%

- 12. The formulation of claim 9 wherein the medicament is a powder25 having a mean particle size of about 1-5 microns.
 - 13. A method for treating mammals comprising administering to said mammals an effective amount the aerosol formulation of claim 1.
- 30 14. A method of treating asthma in mammals comprising administering to a mammal in need of such treatment an effective amount of aerosol formulation consisting essentially of:
- A. a medicament selected from the group comprising 35 albuterol, mometasone furoate, beclomethasone dipropionate, and salts and clathrates thereof;

B. 1.1.1.2.3.3,3 hep	tafluoropropane;
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Optionally an excipient selected from the group consisting of:

propylene glycol diesters of medium chain fatty acids;

triglyceride esters of medium chain fatty acids;

perfluorodimethylcyclobutane;

perfluorocyclobutane;

polyethylene glycol;

10 menthol;

iauroglycol;

diethylalycol monoethylether;

polyglycolized glycerides of medium chain

fatty acids;

15 alcohols;

short chain fatty acids;

eucalyptus oil; and combinations thereof;

optionally a surfactant selected from the group consisting of:

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oleic acid;

sorbitan trioleate;

cetyl pyridinium chloride;

soya lecithin;

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polyoxyethylene (20) sorbitan monolaurate;

polyoxyethylene (20) sorbitan monostearate; polyoxyethylene (20) sorbitan monooleate;

polyoxyethylene (10) stearyl ether;

polyoxyethylene (2) oleyl ether;

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polyoxyethylene-polyoxypropylene-ethylenediamine

block copolymers;

polyoxypropylene-polyoxyethylene block copolymers; castor oil ethoxylate; and combinations thereof; and

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E. optionally one or more additives selected from at least one of the following classes:

preservatives;

buffers;

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antioxidants;

sweeteners; and

taste masking agents.

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I. CLASSIFICATION OF SUI	SJECT MATTER (if several classification	international Applica No PCT	/US 92/04619
	ent Classification (IPC) or to both Natio A 61 K 9/72		
U. FIELDS SEARCHED			
	Minimum D	ocumentation Searched?	
Classification System		Classification Symbols	
Int.Cl.5	A 61 K	C 07 K	
	Documentation Searched to the Extent that such Docum	other than Minimum Documentation nents are Included in the Fields Searched ⁸	
III. DOCUMENTS CONSIDER	RED TO BE RELEVANT ⁹		
	Document, 11 with indication, where app	propriate, of the relevant passages 12	Relevant to Claim No.
			venerant in Claim Mo.
MANUF line	9200062 (MINNESOTA M ACTURING CO.) 9 Janua 1 – page 6, line 39; 13,14, examples 59–6	ry 1992, see page 1, page 11, example 47:	1,2,4-6 ,9-14
1992,	see page 1, line 1 -	200061 (FISONS PLC) 9 January see page 1, line 1 - page 8, line 2; pages examples 4-6,10-16; claims	
MANUF. line	9114422 (MINNESOTA M ACTURING CO.) 3 Octob 1 - page 7, line 12; cclaims	er 1991, see page 1	1,4-6,9 -14
INT. (P111496 (BOEHRINGER GmbH) 8 August 1991, rticular page 5, exam	see the whole document.	1-6,8- 14
Considerat to be of parts earlier document but pub filing date "L" document which may thr which is cited to establish citation or other special r "O" document referring to an other means	neral state of the art which is not niar relevance lished on or after the international w doubts on priority claim(s) or the publication date of another eason (as specified) oral disclosure, use, exhibition or to the international filling date but e claimed	"T" later document published after the interna or priority date and not in conflict with it cited to understand the principle or theor, invention "X" document of particular relevance; the clair cannot be considered novel or cannot be c involve an inventive step "Y" document of particular relevance; the clair cannot be considered to invelve an inventi document is combined with one or more o ments, such combination being obvious to in the art. "&" document member of the same patent fam Date of Mailing of this International Sear	ne application are underlying the med invention considered to med Invention. We step when the ther such docu-a person skilled
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EUROPE	AN PATENT OFFICE	DOGWQF460	xiz

Page 2
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	International Applic No PCT/	us 92/04619
II. DOCUMEN	TS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	Relevant to Claim No.
Category ^o	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to comment
, x	WO,A,9111495 (BOEHRINGER INGELHEIM INT. GmbH) 8 August 1991, see the whole document, in particular page 6, example 5	1-6,8- 14
,х	WO,A,9111173 (FISONS PLC) 8 August 1991, see page 1, line 1 - page 9, line 10; pages 10,11, examples 2,4,6,8; claims	1-14
	100 come short (famoury 1985)	

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9204619

SA 60806

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 01/09/92. The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
WO-A- 9200062		AU-A-	8050891	23-01-92
WO-A- 9200061	09-01-92	AU-A-	8055691	23-01-92
WO-A- 9114422	03-10-91	AU-A- US-A-		21-10-91 02-06-92
WO-A- 9111496	08-08-91	DE-A- AU-A-	4003270 7211691	08-08-91 21 - 08-91
WO-A- 9111495	08-08-91	DE-A-	4003272 7211391	08-08-91 21-08-91
WO-A- 9111173	08-08-91	None		
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